



Subject ID: VM (supplied by FFF Enter)		Subject Initials: First Middle Last									
business hours and on	weekends, ple	at 800-843-7477 to assure an immediate response. After ease select the "emergency order" option.									
After placing your reque it to FFF at 951-296-25		mplete all pages of the downloadable release form and fax									
PLEASE NOTE											
This product is made is required.	de available ir	n the US under BB-IND 7201 reviewed by FDA. IRB review									
▶ Does your organization have a local IRB? ☐ Yes ☐ No											
> FFF requests that local IRBs waive any fees associated for their review of this study.											
The FDA has approved cost recovery for Varicella Zoster IgG under this protocol at \$128.34 per 125 IU vial.											
Subject Information											
Date of birth		MM DD YYYY									
Gender		☐ male ☐ female									
Subject weight		LBS lbs / 2.2 = kilograms									
	KG (required to calculate # of vials)										
Dose											
 125 IU/10 kg IM to a maximum dose of 625 IU (5 vials). Minimum dose is 125 IU (one vial) for patients ≤ 10 kg. 											
	mber of vials required:										
Subject Exposure to Var											
(VariZIG administration	must be withii	n 10 days of exposure to VZV)									
Description of exposure											
Date of first exposure to person infected with VZV											
Time since exposure		DAYS HOURS MINUTES									
Date of appearance of	esions on	■ ■ Not on the black in									
mother (for babies with in-utero exposure only)		M M D D Y Y Y Y									





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(sı	upplied by FFF Enterprises)	irst Midd	le Las	st
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INC	LUSION CRITERIA			
Var	riZIG administration must be within 10 days of exposure to VZV			
	he Subject any of the following at risk patients?			
Che	eck all that apply:		Yes	No
•	Immunocompromised child with no history or evidence of prior infection		Щ	Ц
•	Newborn of mother with VZV < 5 days before or < 2 days after delivery		Щ	닏
•	Premature infant		Щ	Ц
•	Full term infant < 1 year of age		Щ	Ш
•	Immunocompromised adult with no history or evidence of prior infection			
•	Healthy adult with no history or evidence of prior VZV infection			
•	Pregnant woman with no history or evidence of prior VZV infection			
EXL	USION CRITERIA			
	ne answer to any question below is "yes," the subject is not eligible to participa	te in this	trial.	
Tim	ne since exposure cannot exceed 10 days.	,	Yes	No
1.	Does subject have a known immunity to VZV, i.e. previous infection or vaccination		162	NO
	(vaccination = 2 doses of varicella vaccine)?			
2.	Does subject have a history of hypersensitivity to blood or blood products			$\overline{}$
	including IV or IM human immunoglobulin preparations?		Ш	Ш
3.	Is the subject hypersensitive to any component of VariZIG™, its diluent or its pactive. Varicella-Zoster Immune Globulin (Human), sodium chloride, sodium phosph			
	glycine, polysorbate 80, latex stopper)?	iate,	П	
4.	Does the subject have a history of selective immunoglobulin A (IgA) deficiency?		Ħ	\Box
5.	Does the subject have evidence of varicella or zoster lesions prior to dosing?		Ħ	Ħ
6.	Is the subject severely thrombocytopenic (platelets < 50 X 10 ⁹ /L)?			
Phys	sician's Eligibility for Clinical Trials			
	ne answer to question 1 or 2 is " yes " or if the answer to question 3 is " no ," the ph	ysician i	s not	
	gible to participate in this trial.			
1.	Have you ever been disbarred from performing a clinical trial?	`	Yes □	No □
2.	Are you an employee of Cangene Corporation, or have you or your institution		ш	ш
۷.	received a significant benefit (such as payment, proprietary interest or equity) from Cangene Corporation?			
3.	Are you a medical doctor currently licensed in the jurisdiction where treatment wi	II		
	take place and licensed to prescribe medicinal products?			
Ic	ertify that all the above information is true and accurate to the best of (Physician signature on next page)	my kno	wlec	lge.





Subject ID: VM	Subject Initials:
(supplied by FFF Enterprises)	First Middle Last
	Date:
Physician's Signature	MM DD YYYY
Print Name of Physician	
Physician Contact Information (p	lease print)
Hospital or medical facility name:	Street address:
	City:
	State: Zip code:
Phone number (include area code):	()
Fax number (include area code):	<u>()</u>
Email address (required):	
Research Coordinator Contact In	formation (please print)
Name:	Phone number: ()
	Fax Number: (
	Email address:
Pharmacy Contact Information	
Name:	Phone number: ()
	Fax Number: ()
	Email address:
	FFF account number:
	DEA Board of Pharmacy number:





Subject ID: VM		Subject Initials:							
(supplied by FFF Enterprises)				Firs	t Middle Last				
Shipping Address (if different from physician contact information)									
Hospital or medical facility name:	Street address:								
	City:								
	State:		_Zip code:						
	,								
Local IRB Contact Information									
Contact name:									
Phone number: ()									
Email address:									
Completed by FFF Enterprises									
Is subject eligible for the study?				ber of vials: _					
Release authorized by:									
Signature									
	Date:								
Print Name		MM	D D	YYYY					

Notes:

- 1. This product is being provided to fill a gap in therapy in the United States.
- 2. FDA has approved cost recovery for Varicella Zoster IgG under this protocol at \$128.34 per 125 IU vial. Each facility will be responsible for submitting the cost recovery amount to FFF within 30 days of the invoice date.
- 3. Varicella Zoster IgG may be used in only one or very few subjects at each institution. IRB review fees would significantly increase the unit cost of therapy beyond the ability of the sponsor and distributor to recover their costs. Cangene and FFF request that local IRBs waive any fees associated with their review of this study.